

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Attorney Docket No.: DEX-0087

Inventors: Recipon et al.

Serial No. 09/705,500

Filing Date: November 3, 2000

Examiner: Canella, Karen A.

Group Art Unit: 1642

Title: A Novel Method of Diagnosing,
Monitoring, Staging, Imaging and
Treating Cancer

Declaration under Rule § 1.131

We, Herve Recipon, Roberto A. Macina, Sei-Yu Chen and
Yongming Sun, hereby declare:

1. We are co-inventors of the above-referenced patent application.

2. As the co-inventors of the above-referenced patent application, we are familiar with the teachings of the above-referenced patent application.

3. The use of Lng108, also known as fy108, as a cancer diagnostic was conceived and reduced to practice in this country at diaDexus, Inc., prior to October 27, 1999 which was located at that time in Santa Clara, California, USA.

4. Specifically, Lng108 relative expression was determined in accordance with our standard Quantitative Polymerase Chain Reaction (QPCR) protocol and outlined in the above-referenced patent application at page 20, line 16 through page 21, line 13; and page 17, line 12 through page 18, line 9 of priority application U.S. Provisional No. 60/163,144, filed on November 4, 1999. These experiments measuring relative levels of Lng108 in cancerous, normal-adjacent, and normal tissues using Polymerase Chain Reaction in real time were performed prior to October 27, 1999.

Real-Time quantitative PCR with fluorescent Taqman probes is a quantitation detection system utilizing the 5'-3' nuclease activity of Taq DNA polymerase. The method uses an internal fluorescent oligonucleotide probe (Taqman) labeled with a 5' reporter dye and a downstream, 3' quencher dye. During PCR, the 5'-3' nuclease activity of Taq DNA polymerase releases the reporter, whose fluorescence can then be detected by the laser detector of the Model 7700 Sequence Detection System (PE Applied Biosystems, Foster City, CA, USA).

Amplification of an endogenous control was used to standardize the amount of sample RNA added to the reaction and normalize the Reverse Transcriptase (RT) efficiency. Either cyclophilin, glyceraldehyde-3-phosphate dehydrogenase (GADPH) or 18S ribosomal RNA (rRNA) was used as this endogenous control. To calculate relative quantitation between all samples studied, the target RNA levels for one sample were used as the basis for comparative results (calibrator). Quantitation relative to the calibrator can be obtained using the standard curve method or the comparative method (User Bulletin #2: ABI PRISM 7700 Sequence Detection System).

The tissue distribution, and the level of the target gene for every example in normal and cancer tissue were determined. Total RNA was extracted from normal tissues, cancer tissues, and from cancers and the corresponding matched adjacent tissues. Subsequently, first strand cDNA was prepared with reverse transcriptase and the polymerase chain reaction was done using primers and Taqman probe specific to each target gene. The results are analyzed using the ABI PRISM 7700 Sequence Detector. The absolute numbers are relative levels of expression of the target gene in a particular tissue compared to the calibrator tissue.

5. Attached are copies of laboratory notebook pages 0034-124, 0034-168 and 0049-048 containing QPCR experimental procedures and results for Lng108, also known as fy108. The actual dates upon which these experiments were performed in this country at diaDexus, Inc., prior to October 27, 1999 which was located at that time in Santa Clara, California, USA, have been redacted from the attached copies. These experiments, all performed prior to October 27, 1999, demonstrated the use of Lng108 as a diagnostic marker for cancer.

We further declare that all statements made herein of our own knowledge are true and that all statements made on information and belief are believed to true, and further, that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

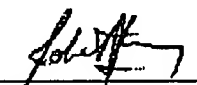


Roberto A. Macina


Date_____
Roberto A. Macina_____
Date_____
Sei-Yu Chen_____
Date_____
Yongming Sun_____
Date

We further declare that all statements made herein of our own knowledge are true and that all statements made on information and belief are believed to true, and further, that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Herve Recipon

Date

Roberto A. Macina

Date

Sei-Yu Chen

Date

Yongming Sun

Date

We further declare that all statements made herein of our own knowledge are true and that all statements made on information and belief are believed to true, and further, that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

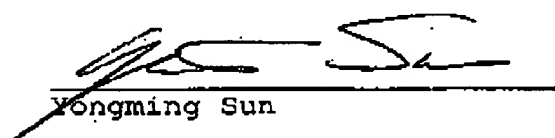
Herve Recipon

Date

Roberto A. Macina

Date

Sei-Yu Chen

Date

Yongming Sun

August 25, 2004
Date

P. 7

$$Fy 108 : N = 20$$

Name James L
 Date _____
 Exp 224 pds = 12 Y
 Cycle 7/804
 (N) 2 Panel Los Mochis

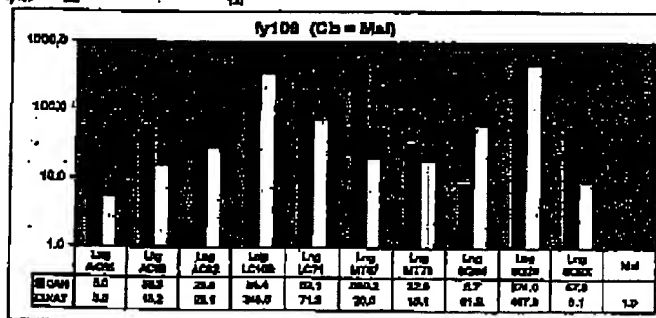
	Water	4.8% (0.9%) DATA
XX	Melting Point	16.0
T Solubility	Soluble in	3.0
T Solubility	Insoluble in	3.0
Flow	Freeze	3.0
	10	10

**Threats (may be repeat threats):**

Matched Tuple Query

[illegible]

TYPE	THROW	CAN	WGT	
1000	Wg AC28	5.0	5.3	
1010	Wg AC38	10.5	10.3	paired Macdonald vs WGT
1020	Wg AC22	20.0	20.1	paired vs 0.1746
1030	Wg LC22	10.1	200.0	
1040	Wg 1571	24.2	24.3	
1050	Wg 1455	24.0	24.0	
1060	Wg MT75	22.0	20.5	
1070	Wg 80M	8.7	41.0	
1080	Wg 8075	27.0	407.0	
1090	Wg 800K	27.0	8.1	
1100		0.0		



WITNESS SIGNATURE:

DATE:

CONTINUED FROM PAGE 167

•N=20 PCR Assay Worksheet

PCR Target LysR
 Tissue Load high
 Endogenous Control 16S rRNA
 Target Primers FIR (anti-Yellow)
 Date of Calibration Mar 7 2004

Name James
 Date 8
 Exp 2240160410
 Order 100/100000
 (Net2 Parcel Lot Number) 2240160410

Master Skills

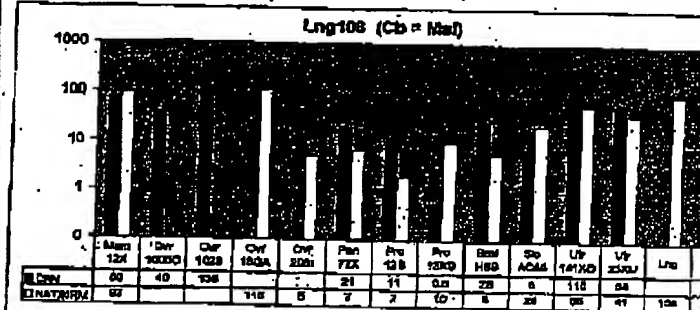
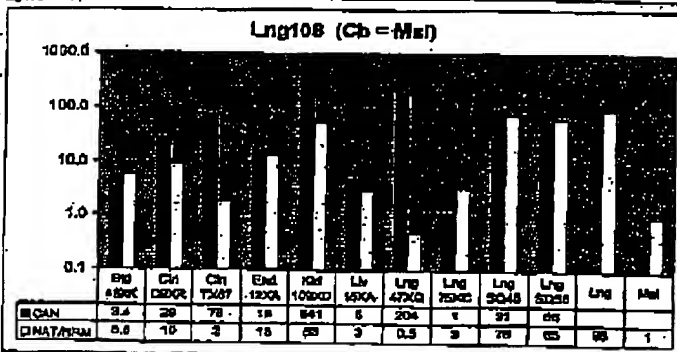
	Wistar	4.88 (20%)	24.1%
22x	Meaningful	15.9	PMLO
12x	Forward	0.08	3.8
100x	Reverse	0.08	3.8
22x	Probe	0.08	3.8

20
10 U. CONA

10 UL CONNA

Timeline (as per report below):

Matched Theses Query

[illegible][illegible]

SCIENTIST SIGNATURE

WITNESS SIGNATURE:

CONTINUED ON PAGE 269

DATE:

DATE:

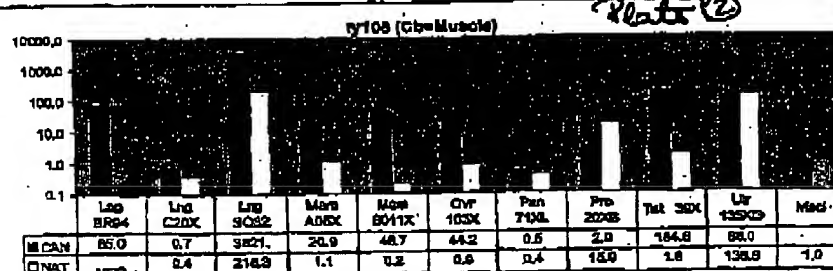
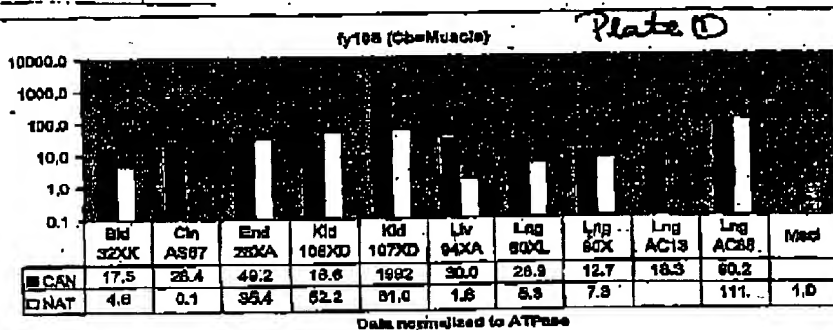
$N=20$ 4108

Objective: to evaluate lung lead levels with additional 400 cases / matched controls.

Protocol:

Timeline (paste report below):

Plots (2) 049, 048B
- same setup as plots (1), but
different trunks.
* - no mycelate added to wells as in
on plots (1)

[illegible]

* Exp. 049 to 048A

	<u>Baseline</u>	<u>Threshold</u>
g108	36-21	0.03
gPAS	36-14	0.02

Exp.	Baseline	Threshold
049	3 to 21	0.02
048B	3 to 14	0.03

CONTINUED ON PAGE 49

SCIENTIST SIGNATURE: *Laura A*

WITNESS SIGNATURE:

DATE:

DATE:

**This Page is Inserted by IFW Indexing and Scanning
Operations and is not part of the Official Record**

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

- ☐ **BLACK BORDERS**
- ☐ **IMAGE CUT OFF AT TOP, BOTTOM OR SIDES**
- ☐ **FADED TEXT OR DRAWING**
- ☐ **BLURRED OR ILLEGIBLE TEXT OR DRAWING**
- ☐ **SKEWED/SLANTED IMAGES**
- ☐ **COLOR OR BLACK AND WHITE PHOTOGRAPHS**
- ☐ **GRAY SCALE DOCUMENTS**
- ☒ **LINES OR MARKS ON ORIGINAL DOCUMENT**
- ☐ **REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY**
- ☐ **OTHER:** _____

IMAGES ARE BEST AVAILABLE COPY.

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.